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TITLE: The BRCA1 Tumor-Suppressor Gene in a Mouse Model of Breast Cancer

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FOREWORD

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In conducting research using animals, the investigator(s) adhered to the "Guide for the Care and Use of Laboratory Animals," prepared by the Committee on Care and use of Laboratory Animals of the Institute of Laboratory Resources, national Research Council (NIE Publication No. 85-23, Revised 1965).
For the protection of human subjects, the investigator(s) adhered to policies of applicable Federal Law 45 CFR 46.
In conducting research utilizing recombinant DNA technology, the investigator(s) adhered to current guidelines promulgated by the National Institutes of Health.
In the conduct of research utilizing recombinant DNA, the investigator(s) adhered to the NIH Guidelines for Research Involving Recombinant DNA Molecules.
In the conduct of research involving hazardous organisms, the investigator(s) adhered to the CDC-NIH Guide for Biosafety in Microbiological and Biomedical Laboratories.

1998 Annual Report for Grant Number DAMD17-96-1-6095

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Appendices: Animal Use Report

USDA Inspection

TABLE OF CONTENTS:

9 10

Page # 1 **Cover Sheet** Standard Form (SF-298) 2 3 Foreword Table of Contents 4 Introduction 5 Body 7 Conclusions 7 References

INTRODUCTION:

In our original application, we proposed to investigate the following specific aims:

1. Characterize of the effects of Brcal expression on the proliferation and differentiation of breast cancer cells of known genotype.

2. Establish mice that lack functional *Brca1* by targeted disruption.

3. Genetic complementation of *Brca1* deficient mice with strains that express oncogenes known to contribute to the development of breast cancer.

To date we have made significant progress on all three aims. In particular, we have studied the effect of overexpressing BRCA1 in mouse mammary epithelial cells and have studied the subcellular localization of the murine Brca1 gene product (Aim 1). These result experiments were completed and summarized in last report. We have created a line of transgenic mice that overexpress human BRCA1 (MBR) (aim 1) and have created a second group of mice that overexpress an antisense construct of mouse Brca1(BAS) (aim 1). We have created a line of mice that carry an inactivated Brca1 locus (BrKO) (aim 2) and have crossed the BrKO mice with mice predisposed to cancer, including (p53^{+/-}, p21^{+/-}, and MMTV-myc) (Aim 3). In the course of these experiments, we experienced several technical difficulties which hindered our progress. In particular, BrKO mice do not develop cancer and mating with cancer prone p53^{+/-}, p21^{+/-}, or MMTV-myc stains did not appear to accelerate or contribute to tumor progression of these strains.

In the last report we proposed substituting BAS (MMTV-Brca1 antisense) mice for BrKO mice in Aim 3 and crossing them into p53^{+/-} and MMTV-myc backgrounds. This experiment is now underway and we have 10-20 animals of each genotype that are being studied for tumor progression. As described below, we have had some indication that the BAS lines are tumor prone indicating that the change in emphasis was justifiable.

BODY:

Analysis of tumor formation in Brca1 knock-out (BrKO) mice:

As described in the 1997 progress report, BrKO mice were established and have been monitored since 1996. In all respects, the phenotype of these mice appears similar to that described by other groups who have knocked out the Brca1 gene (6, 8, 11, 12). Homozygous BrKO mice (BrKO/BrKO) display embryonic lethality at day 6-7.5 of development and thus are not useful for analysis of tumor progression. We also fail to identify tumors in heterozygous animals (BrKO+/-), a result consistent with other groups. To date, we have analyzed over 120 mice that reached 8 mo of age or greater.

In the past year, we have focused on crossing BrKO mice with p53 knockout mice to obtain double heterozygotes. These mice usually develop lymphomas by 6 months of age and are otherwise similar to p53 knockout mice. However, we have had two of these mice develop breast tumors (n=25) and are in the process of breading more animals to verify the phenotype. We have saved DNA from the tumors and will analyze the remaining Brca1 allele for loss. We have plans to submit a request for additional funding to see if irradiation induced DNA damage could accelerate this process.

Generation of Transgenic mice expressing MMTV-Brca1 Antisense and MMTV-BRCA1:

A) To overcome the long latency in tumor progression in BrKO^{+/-}, we generated several lines of transgenic mice expression a Brca1 antisense (BAS) construct targeted to the mammary gland (Table #1). The justification and strategy for creating these mice was described in the 1997 progress report.

The new lines that were created are described in Table 1.

Table 1 Transgenic mice (new lines)

Construct/transgene	Name 1	# of lines1	tumor formation
MMTV-Brca1 antisense	BAS	8	yes (3/8 lines)
MMTV-BRCA1	MBR	1	no
beta-actin-BRCA1		0	N.A. ⁴

¹Number of lines represents the number of founders that transmitted transgenic DNA to offspring.

⁴Not applicable, this construct appears to result in embryonic lethality.

BAS mice have been in the lab for 2.5 years. Upon dexamethasone treatment, 3 week old BAS females show evidence of mammary hyperplasia with increased numbers of branch points in the mammary tree (Figure 1). Non-treated (dexamethasone free) females develop mammary adenocarcinomas with long latency (6-12 months (Figure 2). The incidence is low (4/20 mice > 8mo of age). We are currently mating BAS mice into the p53 null background, following the same procedure outlined for the BrKO mice in the original application. The bigenic BAS/p53 colony is still young, but we have observed 4 mammary tumors thus far and are awaiting additional results as the colony ages. The strategy of targeting Brca1 antisense appears to be a more effective means of eliminating Brca1 expression than waiting for allelic loss at the endogenous Brca1 locus. It therefore is a very promising alternative for studying BRCA1 induced tumors.

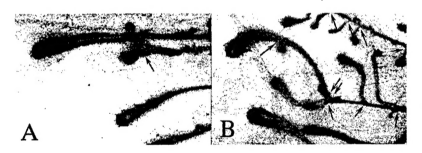


Figure 1 Expression of Brca1 antisense in mammary epithelial cells leads to increased branching and hyperplasia. Panel A shows a whole mount preparation a wild-type mammary gland from a 4 week old mouse treated for 7 days with dexamethasone. Panel B shows a similar preparation from a BAS transgenic litermate. Arrows identify branch points which are much more prevalent in the BAS animal.

We have also generated a number of BASxMMTV-myc mice. At present the females have not developed tumors (average age 3 months) but they are being monitored closely.

Strategy:

The strategy is thus to continue analyzing BrKOxp53 and BAS mice for tumor progression. We are establishing cell lines from the BAS mice in the hopes of demonstrating reduced Brca1 protein expression. We will analyzed all tumors for LOH at the BRCA1 and p53 loci.



Figure 2 Expression of Brcal antisense in mammary epithelial cells leads to the development of mammary adenocarcinomas. The Panel shows a 5 um histological section of mammary tissue obtained from an BAS female. The right side of the panel shows non transformed mammary tissue including ducts (arrows) and adipocytes (a). The left side of the panel shows a well developed adenocarcenoma.

Conclusions:

At the end of year 2, we have made progress on all 3 Aims. We have shown that mouse Brca1 is a nuclear protein that blocks cell proliferation when overexpressed. We have also shown that Brca1 is an essential gene. Loss of the gene in BrKO mice results in early embryonic lethality. Heterozygous BrKO animals are healthy and do not appear to show increased susceptibility to breast cancers, or to any other disease states. While the lack of disease in BrKO mice has been disappointing, the Brca1 antisense (BAS) approach appears to be working. Specifically, we appear to be able to reduce Brca1 protein levels to the point were we can observe increased proliferation (hyperplasias) without inducing cellular lethality. We will continue characterizing these mice. To date 3 out of 8 BAS lines have developed at least one mammary tumor and we are particularly focused on line G which appears particularly cancer prone. We will continue analyzing dexamethasone responsiveness, and sensitivity to DNA damage mediated by ionizing radiation. We will present these results in the final report. As stated, we are in the proceeding with complementation experiments originally proposed in Aim 3 with the BAS transgenics instead of the BrKO mice.

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Definitions of Column	Headings on Back of Fo	orm		······································	
A. Animal	B. Number of animals purchased, bred, or housed but not yet used	C. Number of animals used involving no pain or distress	D. Number of animals used in which appropriate anesthetic, analgesic, or tranquilizing drugs were used to alleviate pain	E. Number of animals used in which pain or distress was not alleviated	F. Total Number of Animals (Columns C+D+E
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Rats					
Fish				·	
List Others:					

^{*}AAALAC - Association for the Assessment and Accreditation of Laboratory Animal Care



United States Department of Agriculture Animal and Plant Health Inspection Service Animal Care

INSPECTION REPORT

University of California-Los Angeles Department of Laboratory Animal Medicine 630 Circle Drive Site 01 Main Vivarium 93-R-0044 AG 05-22,27-98 0930 Routine

NARRATIVE

Animal Inventory- 18 Dogs

Los Angeles, CA

14 Cats

35 Guinea pigs

11 Hamsters

208 Rabbits

21 Chinchillas

38 Deer mice

25 Non-human primates(11 baboons, 2 macaques, 12 squirrel monkeys)

46 Domestic swine

CATEGORY III: Non-compliant item(s) identified this inspection

Section 3.2(d), 3.26(d), 3.51(d), 3.75(c)-<u>Interior Surfaces</u> - The following areas were noted with unsealed, porous surfaces which can not be cleaned and sanitized as required-

- peeling paint, pitted floors, wallboards separating from walls (porous

wallboard and concrete exposed) - rooms housing non-human primates

- tile missing from wall (porous wallboard exposed)- guinea pig room

- brick floors with peeling sealant- rabbit room

- peeling paint on walls- rooms housing dogs

- peeling paint, pitted floors (porous wallboard and concrete)- cagewashing and bedding storage rooms

All above to be corrected by: 7-3-98

Note: All above reported to facility maintenance department prior to this inspection.

Prepared By: 16 thee m to low DVm vnc	Date: 1-3-98
Title: Kathleen M. Garland, Veterinary Medical Officer, USDA, APHIS, Animal Care	LARIS 10 NO. 5006
Copy Received By: 3. 16	Date: 6/3/88
Title:	

Section 3.11(c), 3.31(b),3.56(c), 3.84(c), 3.131(c)-Housekeeping - Floor drain of from dog recovery room- Florescent light not functioning properly in surgery of the control of the cont	cover missing of Josephania com- Te-bo
from dog recovery room— Florescent light not functioning properly in surgery of the corrected by: Corrected by: Corrected by: Corrected by: Corrected by: Corrected by: Wall ventilation duct cover noted with build-up of discovery noted by:	rt and dust in
surgery room- Corrected at time of inspection	
Section 3.129(b)-Feeding - Pigs are fed by placing feed in rubber tub. Section 2.33(b)-Veterinary Care - rubber tubs noted with areas missing (che areas that have been worn and chewed so that fibers are exposed- these tubs cleaned and sanitized as required. Also, there is a concern that the animals may pieces or fibers of rubber which may result in impaction or other injury to the animals represented by: To be corrected by:	an not be kept be ingesting mals.
Serviced medications noted mixed with curre	nt dated drugs in
Depoprovera- expired 2-97 Oxytetracycline-expired 7-97	
Heparin-expired April 1,1998	
	ion
All above were removed at time of inspection - corrected at time of inspect	
Section 3.81- Environmental Enhancement - This facility houses baboons, send macaques, but does not account for, or document, species differences in program of environmental enhancement/ behavioral enrichment. Although it is written plan that each non-human primate room is provided with intercom-supp throughout the day, it was not noted at time of inspection. To be corrected by: Note: Development of environmental enhancement plans, protocol literature protocol / program reviews were discussed.	lied music 7-3-98
End of Report	
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Prepared By: 16 H. Gariand, Veterinary Medical Officer, USDA, APHIS, Animal Care	LARIS ID NO. 5006
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